=> file reg
FILE 'REGISTRY' ENTERED AT 12:51:21 ON 03 JUN 2004
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=> file hcaplus FILE 'HCAPLUS' ENTERED AT 12:51:25 ON 03 JUN 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 3 Jun 2004 VOL 140 ISS 23 FILE LAST UPDATED: 2 Jun 2004 (20040602/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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		E T	RISULFATE"/CN		-, -, -, -, -, -, -, -, -, -, -, -, -, -
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L34	108	SEA	FILE=REGISTRY	ABB=ON	C19H24N2.CLH/MF
L35	17	SEA	FILE=REGISTRY	ABB=ON	L25 AND L34
L38	1	SEA	FILE=REGISTRY	ABB=ON	L35 AND DIMETHYLAMINOPROP?
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L58	1	SEA	FILE=REGISTRY	ABB=ON	L57 AND PYRIMI?
L59	1	SEA	FILE=REGISTRY	ABB=ON	C4H6N4S.H2O4S/MF

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L62
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L63
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L64
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                                         L62 (L) ELECTROLYTE?
1.65
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L79
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L81
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                                          "1,4,7,10,13,16-HEXAAZACYCLOOCTADECAN
                E TRISULFATE"/CN
L91
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L92
           1122 SEA FILE=HCAPLUS ABB=ON L91
T.94
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L95
              6 SEA FILE=HCAPLUS ABB=ON L92(L)ELECTROLYT?
L96
              8 SEA FILE=HCAPLUS ABB=ON
                                         L65 OR L94 OR L95
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L95 ANSWER 1 OF 6 HCAPLUS COPYRIGHT 2004 ACS on STN
     2001:731243 HCAPLUS
DN
     135:291346
TΙ
     Secondary lithium batteries
     Yang, Li; Yoshida, Toshihiro; Nemoto, Hiroshi; Takahashi, Michio
     NGK Insulators, Ltd., Japan
SO
     PCT Int. Appl., 67 pp.
                                             applicant
     CODEN: PIXXD2
DΤ
     Patent
LA
     Japanese
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
                                                            DATE
                                           ------
PΤ
     WO 2001073884
                      A1
                            20011004
                                           WO 2001-JP1135
                                                            20010216
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             PT, SE, TR
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                      A2
                            20011005
                                           JP 2000-89934
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     JP 2001283907
                      A2
                            20011012
                                           JP 2000-89936
                                                            20000328
                      A2 '
     JP 2001283919
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                                           JP 2000-89965
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     JP 2001283920
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     JP 2001283921
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    US 2003190530
                            20031009
                      A1
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                      Α
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                      Α
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                      Α
                            20000328
    JP 2000-89974
                      Α
                            20000328
    WO 2001-JP1135
                      W
                            20010216
AΒ
    The batteries have a coiled electrode/separator stack and a nonaq. Li salt
    electrolyte solution, where the cathode, anode, separator, and/or the
```

electrolyte solution contain organic and/or inorg. Cu corrosion inhibitor or Cu trapping agent, a compound containing both basic organic groups and inorg. acid groups, a N-O radical containing cyclic compound, a compound not containing Lewis acid

atoms and Lewis base atoms at the same time, a 3-dimensional siloxane compound, and/or a nonionic surfactant, and/or a cyclic Mn2+ source in the electrolytes; and the electrolyte soln contains a water trapping agent or a HF trapping agent.

ICM H01M010-40 IC

CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology)

ΙT 95-14-7, 1,2,3-Benzotriazole 128-94-9, 1,8-Diamino-4,5dihydroxyanthraquinone 2564-83-2 9004-99-3 9014-92-0, Polyethylene glycol mono-dodecylphenyl ether 14325-24-7, Manganese (II) phthalocyanine 14691-88-4 16011-96-4, 2-Iminopiperidine hydrochloride 26027-38-3, Polyethylene glycol mono-4-nonylphenyl ether 26635-92-7 34272-83-8 207505-82-6 213453-16-8 364589-08-2 364589-09-3

RL: MOA (Modifier or additive use); USES (Uses) (additives for electrodes and separators and electrolyte solns. in secondary lithium batteries)

16011-96-4, 2-Iminopiperidine hydrochloride IT RL: MOA (Modifier or additive use); USES (Uses)

(additives for electrodes and separators and electrolyte solns. in secondary lithium batteries)

RN 16011-96-4 HCAPLUS

CN 2-Pyridinamine, 3,4,5,6-tetrahydro-, monohydrochloride (9CI) (CA INDEX NAME)

HC1

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L95 ANSWER 2 OF 6 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1987:55762 HCAPLUS

DN 106:55762

TI Total intensity and quasielastic light scattering studies on the association of amphiphilic drugs in aqueous electrolyte solutions

UΑ Attwood, D.; Fletcher, P.

CS Dep. Pharm., Univ. Manchester, Manchester, M13 9PL, UK

SO Journal of Colloid and Interface Science (1987), 115(1), 104-9 CODEN: JCISA5: ISSN: 0021-9797

DT Journal

LA English

Total intensity and quasielastic light scattering (QELS) techniques were AB used to examine the effect of electrolyte on the micellar properties of the antidepressant drug imipramine-HCl [113-52-0] in aqueous solution A progressive increase of aggregation number and micellar radius was observed with increase of NaCl concentration over the range 0.05 to 0.6M.

A marked decrease of the diffusion coefficient with increase of $\ensuremath{\operatorname{drug}}$ concentration at

concns. just in excess of the critical micelle concentration was noted and discussed. QELS measurements of trimetaphan camphorsulfonate [68-91-7], which exhibits continuous association, showed a complex variation of effective diffusion coefficient with drug and electrolyte concentration

CC 63-5 (Pharmaceuticals)

IT 68-91-7 113-52-0, Imipramine hydrochloride

RL: PRP (Properties)

(association of, in **electrolyte** solns., total intensity and quasielastic light scattering in study of)

IT 113-52-0, Imipramine hydrochloride

RL: PRP (Properties)

(association of, in **electrolyte** solns., total intensity and quasielastic light scattering in study of)

RN 113-52-0 HCAPLUS

5H-Dibenz[b,f]azepine-5-propanamine, 10,11-dihydro-N,N-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

CN

● HCl

L95 ANSWER 3 OF 6 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1986:135972 HCAPLUS

DN 104:135972

- TI Adsorption of antidepressants on clays. Part II. Effect of solvents and electrolytes on the adsorption of imipramine hydrochloride on veegum and bentonite
- AU El-Mowafi, Mohamed A.; Geneidi, Ahmed S.; Kassem, Aly A.

CS Nile Co. Pharm., Cairo, Egypt

Ι

SO Egyptian Journal of Pharmaceutical Sciences (1985), Volume Date 1983, 24(1-4), 287-96 CODEN: EJPSBZ; ISSN: 0301-5068

DT Journal

LA English

GI

AB Increasing concns. of polyethylene glycol 400 [25322-68-3], EtOH [64-17-5] and glycerol [56-81-5] decreased the adsorption of imipramine-HC1 (I-HC1) [113-52-0] on Veegum S-6198 [1327-43-1] and bentonite, while sorbitol [50-70-4] showed no such effect. With increasing concns. of NaZHPO4, NaCl and Na citrate [68-04-2] the amount of I adsorbed increased. Reasons for the difference in behavior are discussed.

CC 63-5 (Pharmaceuticals)

IT 113-52-0

RL: PEP (Physical, engineering or chemical process); PROC (Process) (adsorption of, on Veegum and bentonite, solvents and electrolytes effect on)

IT 113-52-0

RL: PEP (Physical, engineering or chemical process); PROC (Process) (adsorption of, on Veegum and bentonite, solvents and electrolytes effect on)

RN 113-52-0 HCAPLUS

CN 5H-Dibenz[b,f]azepine-5-propanamine, 10,11-dihydro-N,N-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

HC1

L95 ANSWER 4 OF 6 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1985:427182 HCAPLUS

DN 103:27182

TI Studies on the adsorption of cationic amphiphilic drugs on antacids and adsorbents. 5. Studies on the desorption behavior of cationic, amphiphilic drugs

AU Thoma, K.; Lieb, H.

CS Inst. Pharm. Lebensmittelchem., Ludwig-Maximilians-Univ. Muenchen, Munich, D-8000/2, Fed. Rep. Ger. SO Pharmaceutica Acta Helvetiae (1985), 60(4), 98-105

CODEN: PAHEAA; ISSN: 0031-6865

DT Journal

LA German

AB The desorption of cationic, amphiphilic drugs such as diphenhydramine-HCl (I) [147-24-0], chlorpromazine-HCl (II) [69-09-0], and imipramine-HCl [113-52-0] from antacids such as Mg trisilicate, basic Bi nitrate and adsorbents such as kaolin was investigated. The desorption was affected very strongly by solvent media, drug and the adsorbent concns. Increasing the elution time increased the desorbed amts. of I from 59.1% to 63.1%. Increasing the electrolyte content and decreasing the pH of the medium increased the desorbed amts. With II, desorption increased as the adsorbent was changed from Mg trisilicate to basic Bi nitrate (maximum).

CC 63-5 (Pharmaceuticals)

ANSWER 5 OF 6 HCAPLUS COPYRIGHT 2004 ACS on STN ΑŃ 1981:20301 HCAPLUS DN 94:20301 ΤI Colloid association of tri- and tetracyclic antidepressives and neuroleptics. 4. Communication. Determination of miceller weights and influence of the medium conditions on the association properties ΑU Thoma, K.; Albert, K. CS Inst. Galenische Pharm., Johann-Wolfgang-Goethe-Univ., Frankfurt, D-6000, Fed. Rep. Ger. SO Pharmaceutica Acta Helvetiae (1980), 55(5), 146-55 CODEN: PAHEAA; ISSN: 0031-6865 יייח Journal LA German AB The micellar wts. and aggregation nos. of 13 tricyclic antidepressants and neuroleptics in electrolyte-containing solns. were determined In acetate buffer pH 5.0, aggregates with micellar wts. between 3900-17,600 were formed. In a physiologic saline solution micellar wts. range from 5400 to 13,100. In 0.9% NaCl solution the piperazine derivs. clopenthixol-2HCl [633-59-0], flupentixol-2HCl [2413-38-9], and thiothixene-2HCl [49746-04-5] have on an average 8 monomers smaller micelles, and aminopropyl derivs. have 3 to 14 monomers larger micelles than in acetate buffer. The decrease and the increase of the size of the aggregates, can be explained in terms of changes in pH and counterions. The effect of different electrolytes on micelle formation of clomipramine-HCl [17321-77-6] and clopenthixol-2 HCl was investigated. In comparison to NaCl the same molar concns. of Na methanesulfonate [2386-57-4], NaOAc [127-09-3] and Na lactate [72-17-3] increase the critical micelle concns. (cmc), whereas Na succinate [150-90-3], Na tartrate [868-18-8], Na maleate [371-47-1], Na benzenesulfonate [515-42-4] and Na citrate [68-04-2] decrease the cmc's. The addition of electrolytes leads to a depression of the cmc's of clomipramine-HCl and clopenthixol-2 HCl. By increasing the concentration of chloride ions from 0.03 to 0.3 mol.1-1, the hydration of the clomipramine-HCl monomers was decreased and the cmc decreases to one third of its original value. With a simultaneous change of the dissociation degree of a compound, the pH can significantly influence the cmc. For the diacidic piperazine derivative clopenthixol-2 HCl the following was shown: increasing the pH from 2 to 6.5 decreased the dissociation degree of the weak basic N and because of the reinforced hydrophobic character of the compound the cmc also decreases to 4.4 10-4 mol.1-1. CC 63-5 (Pharmaceuticals) ΙT 58-28-6 **113-52-0** 549-18-8 633-59-0 739-71-9 894-71-3 1225-55-4 1229-29-4 2413-38-9 6469-93-8 10563-70-9 17321-77-6 49746-04-5 RL: BIOL (Biological study) (aggregation and micellar wts. of, electrolytes and pH effect on) IT 113-52-0 RL: BIOL (Biological study) (aggregation and micellar wts. of, electrolytes and pH effect on)

5H-Dibenz[b,f]azepine-5-propanamine, 10,11-dihydro-N,N-dimethyl-,

monohydrochloride (9CI) (CA INDEX NAME)

113-52-0 HCAPLUS

RN CN

HC1

L95 ANSWER 6 OF 6 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1970:402473 HCAPLUS

DN 73:2473

TI Cerebrospinal fluid electrolyte changes during lateral ventricular perfusion with noradrenaline and associated cardiovascular responses

AU Melville, Kenneth I.; Johnson, Melvin C.
CS Dep. Pharmacol. Ther., McGill Univ., Montreal, QC, Can.

SO Neuropharmacology (1970), 9(2), 79-95

CODEN: NEPHBW; ISSN: 0028-3908

DT Journal LA English

AB Changes of the concentration of Na+, K+, Ca2+, and Mg2+ in cerebrospinal fluid, blood pressure, and heart rate elicited by a 60-min perfusion of the lateral ventricle were studied in chloralose-anesthetized, vagotomized cats. Perfusion with artificial cerebrospinal fluid changed neither the electrolytes of the cerebrospinal fluid nor the blood pressure of heart rate. Perfusions with artificial cerebrospinal fluid containing 1-noradrenaline bitartrate (0.08-8 μg/ml) decreased Na+ concns. in the cerebrospinal fluid and elicited hypotension wihout changing the heart rate. Perfusions with high doses of noradrenaline (80-160 μ g/ml) increased K concns. in the cerebrospinal fluid and elicited hypertension and tachycardia. With prolonged perfusions, tolerance or tachyphylaxis developed to these responses to noradrenaline. Intraventricular pretreatment with reserpine (1 mg), tranylcypromine sulfate (5 mg), chlorpromazine-hCl (10 mg), or imipramine-HCi (2 mg) increased K+ and Mg2+ concns. in the cerebrospinal fluid. Ca2+ concns. were also increased by chlorpromazine and imipramine, but Na+ concns. were not affected. After reserpine or tranylcypromine pretreatment, the increase of K+ concentration,

hypertension, and the tachycardia induced by noradrenaline were antagonized. After chlorpromazine or imipramine pretreatment, noradrenaline still induced hypertension and tachycardia, associated with no further change in K+ but with a decrased in Mg2+ concns. Centrally mediated cardiovascular responses induced by intraventricular noradrenaline seem to involve changes in cerebrospinal fluid and possibly in brain Na+ and (or) K+, and reserpine, tranylcypromine, chlorpromazine, and imipramine appear to induce significant shifts in brain Mg2+ and ca2+ concns. Which might explain some actions of these drugs on the central nervous system.

CC 15 (Pharmacodynamics)

IT 50-55-5, biological studies 51-40-1 69-09-0 **113-52-0** 13492-01-8

RL: BIOL (Biological study)

(electrolytes of cerebrospinal fluid after administration of)

● HCl

=> =>	d que	
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L35		SEA FILE=REGISTRY ABB=ON L25 AND L34
L38		SEA FILE=REGISTRY ABB=ON L35 AND DIMETHYLAMINOPROP?
L53		SEA FILE=REGISTRY ABB=ON C4H7N5O.H2O4S/MF
L57		SEA FILE=REGISTRY ABB=ON C4H9N3.2CLH/MF
L58		SEA FILE=REGISTRY ABB=ON L57 AND PYRIMI?
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		L20
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L64		SEA FILE=HCAPLUS ABB=ON L62(L)ELECTROLYTE?
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L66		SEA FILE=HCAPLUS ABB=ON (LI OR LITHIUM) (5A) BATTER?
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	_	SULFATE"/CN
L81	1	SEA FILE=REGISTRY ABB=ON "2-IMINOPIPERIDINE HYDROCHLORIDE"/CN
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1.97	ANSWER 1 OF	7	HCAPLUS COD	VRICUT	2004 ACS on STN

ANSWER 1 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 2002:886243 HCAPLUS

DN 137:387083

ED Entered STN: 22 Nov 2002

Nonaqueous gel composition containing crosslinked polymer having ΤI alkylammonium or piperazinium structure and electrochemical cell

ΤN Aizawa, Wakana; Ikegami, Koshiro; Takada, Masakazu; Takaoka, Kazuchiyo

PA Mitsubishi Paper Mills, Ltd., Japan; Nippon Unicar Co., Ltd.

SO Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

IC ICM C08L101-02

ICS H01B001-06; H01G009-025; H01M010-40

CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology)

Section cross-reference(s): 38, 76

FAN.CNT 1

PATENT NO	. KIND	DATE	APPLICATION NO.	DATE
PI JP 200233 PRAI JP 2001-1		20021122 20010509	JP 2001-138273	20010509
GI				

AB The title gel composition comprises a polymer having a crosslinked structure R1NX, R2NYNX2, R4NYNX, I, or II [R1-R7 = (substituted) C1-9 alkyl; X =monovalent inorg. or organic acid or its

equivalent; Y = C1-8 alkylene, alkylene oxide, or xylylene]. The composition, especially

suitable for secondary Li batteries and capacitors,

has high resistance to free acids generated in an electrolyte solution

ST alkylammonium crosslinked polymer electrolyte capacitor; piperazinium crosslinked polymer nonag electrolyte battery

TΤ Capacitors

(double layer; nonaq. gel electrolyte composition containing crosslinked polymer

having alkylammonium or piperazinium structure for battery and capacitor)

IT Secondary batteries

(lithium; nonaq. gel electrolyte composition containing crosslinked polymer having alkylammonium or piperazinium structure for battery and capacitor)

IT Battery electrolytes Crosslinking agents Electrolytic capacitors Polymer electrolytes

(nonaq. gel electrolyte composition containing crosslinked polymer having alkylammonium or piperazinium structure for battery and capacitor)

IT Quaternary ammonium compounds, uses RL: DEV (Device component use); PNU (Preparation, unclassified); PREP (Preparation); USES (Uses) (polymers; nonag, gel electrolyte composition containing crosslinked polymer

having alkylammonium or piperazinium structure for battery and capacitor)

TΤ 96-49-1, Ethylene carbonate 105-58-8, Diethyl carbonate 108-32-7. Propylene carbonate 429-06-1, Tetraethylammonium tetrafluoroborate 21324-40-3, Lithium hexafluorophosphate RL: DEV (Device component use); USES (Uses)

(composition containing; nonag, gel electrolyte composition containing crosslinked

polymer having alkylammonium or piperazinium structure for battery and capacitor) IΤ 1072-63-5DP, N-Vinylimidazole, polymers with alkylammonium compound and piperazinium compound 1337-81-1DP, Vinylpyridine, polymers with vinyl monomer and piperazinium compound 52352-11-1DP, Vinylbenzylamine,

alkylammonium fluorophosphates, polymers with piperazinium compound 476013-44-2P 476013-47-5P 476013-48-6DP, polymers with vinyl monomer and piperazinium compound 476013-50-0DP, polymers with vinyl monomer RL: DEV (Device component use); PNU (Preparation, unclassified); PREP (Preparation); USES (Uses)

- (nonag. gel electrolyte composition containing crosslinked polymer having alkylammonium or piperazinium structure for battery and capacitor)
- L97 ANSWER 2 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN
- AN 2002:80804 HCAPLUS
- DN 136:388363
- ED Entered STN: 31 Jan 2002
- ΤI New technique for preparation of the cathode material of lithium -ion battery
- Wei, Jin-Ping; Deng, Bin; Yan, Jie; Zhou, Zhen; Song, De-Ying ΑU
- CS Institute of New Energy Material Chemistry, Nankai University, Tianjin, 300071, Peop. Rep. China
- SO Gaodeng Xuexiao Huaxue Xuebao (2001), 22(10, Suppl.), 9-12 CODEN: KTHPDM; ISSN: 0251-0790
- PB Gaodeng Jiaoyu Chubanshe
- DTJournal
- LA Chinese
- CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
- The cathode material LiCoO2 of lithium-ion batteries can be obtained from the reaction of LiOH-H2O and Co(OH)2-2xRx(R is the inorg. or organic acid radicals), which is the midst product of the electrolysis of the metal cobalt in a given electrolyte. The achieved cathode material LiCoO2 for lithium -ion batteries has a single phase and its granularity distribution relatively concs. through the expts. of XRD, SEM and Laser Granularity Testing. The discharge capacity is relatively high which reaches 146 mAh/g at the first time of discharging and still maintains 142 mAh/g after 10 cycles of charge/discharge expts. The cost for the preparation of the cathode material LiCoO2 can greatly be decreased through this technique, which has a well-promising application future.
- ST battery cathode cobalt lithium oxide

TT Battery cathodes (new technique for preparation of the cathode material of lithium -ion battery) ΙT 12190-79-3P, Cobalt lithium oxide LiCoO2 RL: PEP (Physical, engineering or chemical process); PNU (Preparation, unclassified); PREP (Preparation); PROC (Process) (new technique for preparation of the cathode material of lithium -ion battery) L97 ANSWER 3 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN AN 2001:865039 HCAPLUS DN 136:9048 ED Entered STN: 30 Nov 2001 Manufacture of cathode active mass and nonaqueous electrolyte battery TΙ using the active mass IN Takagi, Shigeharu PA Toyota Motor Corp., Japan Jpn. Kokai Tokkyo Koho, 7 pp. SO CODEN: JKXXAF DT Patent LA Japanese IC ICM H01M004-58 ICS H01M010-40 CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology) FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE ____ PΙ JP 2001332261 A2 20011130 JP 2000-150317 20000522 PRAI JP 2000-150317 20000522 The cathode active mass is prepared by soaking a Li containing multiple oxide, having a rock salt structure, in water, polar organic solvent, and/or inorg. acid for ≥12 h. The battery is a secondary Li battery. ST secondary lithium battery cathode active mass manuf; lithium multiple oxide manuf secondary battery cathode TΤ Acids, uses RL: NUU (Other use, unclassified); USES (Uses) (inorg., polar; solution soaking of lithium containing multiple oxides in manufacture of cathode active mass for secondary lithium batteries) TΤ Solvents (organic, polar; solution soaking of lithium containing multiple oxides in manufacture of cathode active mass for secondary lithium batteries) ΙT Battery cathodes (solution soaking of lithium containing multiple oxides in manufacture of cathode active mass for secondary lithium batteries TΤ 193214-24-3P, Aluminum cobalt lithium nickel oxide (Al0.05Co0.15LiNi0.802) RL: DEV (Device component use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses) (manufacture of cathode active mass by solution soaking of lithium containing multiple oxides for secondary lithium batteries) 7732-18-5, Water, uses TΤ RL: NUU (Other use, unclassified); USES (Uses) (solution soaking of lithium containing multiple oxides in manufacture of cathode active mass for secondary lithium batteries)

```
L97 ANSWER 4 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN
    1999:212891 HCAPLUS
AN
DN
    130:225376
ED
     Entered STN: 05 Apr 1999
     Battery cathodes and their manufacture
TΙ
     Morita, Kimihiro; Shimizu, Hiroyuki
IN
     Asahi Chemical Industry Co., Ltd., Japan
PΑ
SO
     Jpn. Kokai Tokkyo Koho, 7 pp.
     CODEN: JKXXAF
DT
     Patent
LA
     Japanese
IC
     ICM H01M004-02
     ICS H01M004-04; H01M004-58; H01M004-62; H01M004-66; H01M010-40
CC
     52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
FAN.CNT 1
     PATENT NO.
                   KIND DATE
                                           APPLICATION NO. DATE
     -----
                                           -----
PI JP 11086846 A2 19990330
PRAI JP 1997~180877 19970707
                                           JP 1997-334099 19971204
                            19970707
     The cathodes have a slurry containing Li Ni oxide, a conductive additive, a
     fluoropolymer, a diluting agent, and a slurry stabilizer applied on an Al
     collector. The stabilizer is an inorg. or organic
     acid, Fe bromide, or FeCl3; the Li Ni oxide may contain other
transition metals and elements selected from B, Al, In, Sn, Mg, and Zn.
     The cathodes are prepared by dissolving the stabilizer and the fluoropolymer
     in the diluting agent, dispersing Li Ni oxide and the conductive additive to
     the solution, applying the slurry to Al collectors, and drying.
ST
     battery lithium nickel oxide cathode manuf
IT
     Battery cathodes
        (compns. and manufacture of lithium nickel oxide based cathodes
        from slurries on aluminum substrates for batteries)
IT
     Fluoropolymers, uses
     RL: MOA (Modifier or additive use); USES (Uses)
        (compns. and manufacture of lithium nickel oxide based cathodes from
        slurries on aluminum substrates for batteries)
IT
     221154-74-1, Cobalt lithium magnesium nickel oxide (Co0.1LiMg0.01Ni0.8902)
     RL: DEV (Device component use); PEP (Physical, engineering or chemical
     process); PROC (Process); USES (Uses)
        (compns. and manufacture of lithium nickel oxide based cathodes from
        slurries on aluminum substrates for batteries)
ΙT
     24937-79-9, Poly(vinylidene fluoride)
     RL: MOA (Modifier or additive use); USES (Uses)
        (compns. and manufacture of lithium nickel oxide based cathodes from
        slurries on aluminum substrates for batteries)
TΤ
     872-50-4, N-Methyl-2-pyrrolidone, uses 7705-08-0, Ferric chloride, uses
     128151-86-0, Dynomil
     RL: NUU (Other use, unclassified); USES (Uses)
        (compns. and manufacture of lithium nickel oxide based cathodes from
        slurries on aluminum substrates for batteries)
L97 ANSWER 5 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN
AN
     1997:491191 HCAPLUS
DN
     127:151031
ED
     Entered STN: 04 Aug 1997
ΤI
    Nonaqueous secondary batteries with anodes containing amorphous chalcogen
     compounds or oxides
IN
    Maekawa, Yukio; Miyaki, Yukio
PA
     Fuji Photo Film Co., Ltd., Japan
```

```
Jpn. Kokai Tokkyo Koho, 11 pp.
     CODEN: JKXXAF
DT
     Patent
LA
     Japanese
IC
     ICM H01M004-58
     ICS H01M004-02; H01M004-04; H01M010-40
CC
     52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
FAN.CNT 1
     PATENT NO.
                    KIND DATE
                                          APPLICATION NO.
     -----
                                          -----
                     A2 19970722
PΙ
     JP 09190819
                                          JP 1996-142
                                                           19960105
PRAI JP 1996-142
                           19960105
     The title batteries use Li-intercalating anodes containing
     amorphous chalcogen compds. or amorphous oxides treated with organic acids or
     their salts. The batteries have long cycle life and high capacity.
ST
     anode amorphous chalcogen compd battery; amorphous oxide anode
     lithium battery; carboxylic acid treatment anode battery
IT
     Carboxylic acids, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (amorphous chalcogen compds. or oxides treatment by; anodes for nonaq.
        batteries with capacity and long cycle life)
IΤ
     Chalcogenides
     Oxides (inorganic), uses
     RL: DEV (Device component use); USES (Uses)
        (amorphous, organic acid-treated, anodes; nonag.
        batteries with capacity and long cycle life)
ΙT
        (anodes containing organic acid-treated amorphous chalcogen compds. or
oxides
        for nonag, batteries)
IT
     Secondary batteries
        (lithium; anodes containing organic acid-treated amorphous chalcogen
        compds. or oxides for nonaq. batteries)
IΤ
     57-11-4, Octadecanoic acid, uses 64-19-7, Acetic acid, uses
     Benzoic acid, uses 75-75-2, Methanesulfonic acid 79-09-4, Propionic
     acid, uses 88-99-3, 1,2-Benzenedicarboxylic acid, uses 103-82-2,
     Phenylacetic acid, uses 110-15-6, Butanedioic acid, uses 110-16-7,
     2-Butenedioic acid (Z)-, uses 110-94-1, Glutaric acid 123-76-2,
     Levulinic acid 124-04-9, Adipic acid, uses 127-17-3, Pyruvic acid,
     uses 141-82-2, Malonic acid, uses 144-62-7, Ethanedioic acid, uses
     298-12-4, Glyoxylic acid 528-44-9, Trimellitic acid 541-50-4,
     Acetoacetic acid, uses 553-91-3, Dilithium oxalate
                                                          2051-95-8,
     3-Benzoylpropionic acid 9003-01-4, Polyacrylic acid 25087-26-7,
     Polymethacrylic acid
     RL: NUU (Other use, unclassified); USES (Uses)
        (amorphous chalcogen compds. or oxides treatment by; anodes for nonag.
        batteries with capacity and long cycle life)
ΤТ
     188198-63-2 193217-90-2 193217-92-4 193217-94-6 193217-95-7
     193266-49-8
     RL: DEV (Device component use); USES (Uses)
        (amorphous, organic acid-treated, anode; nonaq. batteries with capacity
        and long cycle life)
IT
     193266-48-7P
     RL: DEV (Device component use); PNU (Preparation, unclassified); PREP
     (Preparation); USES (Uses)
        (amorphous, organic acid-treated, anode; nonaq. batteries with capacity
        and long cycle life)
L97 ANSWER 6 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN
```

AN 1980:498508 HCAPLUS

DN 93:98508

ED Entered STN: 12 May 1984

TI Manganese dioxide, and electrochemical cell having cathode containing said manganese dioxide

IN Hunter, James Charles

Union Carbide Corp., USA PA

SO Eur. Pat. Appl., 20 pp.

CODEN: EPXXDW DT Patent

LA English

FAN. CNT 1

C01G045-02; H01M004-50 IC

52-2 (Electrochemical, Radiational, and Thermal Energy Technology)

Section cross-reference(s): 49

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΙ	EP 9934	A1	19800416	EP 1979-302024	19790928
	EP 9934	B1	19821103		
	R: DE, FR,	GB			
	US 4246253	A	19810120	US 1978-947120	19780929
	CA 1134595	A1	19821102	CA 1979-335495	19790912
	AU 7951218	A1	19800403	AU 1979-51218	19790926
	NO 7903102	A	19800401	NO 1979-3102	19790927
	BR 7906195	A	19800909	BR 1979-6195	19790927
	ES 484508	A1	19801101	ES 1979-484508	19790927
	DK 7904086	A	19800330	DK 1979-4086	19790928
	JP 55100224	A2	19800731	JP 1979-125241	19790928

19830726

19810901

19820126

PRAI US 1978-947120 19780929 MnO2 useful in the manufacture of battery cathodes is prepared by acid

of LiMn2O4. Thus, the preparation of λ -MnO2 by treating LiMn2O4 with 3N HCl, 4.7N H2SO4, or 4N HNO3 is reported. LiMn2O4 was prepared by heating a powdered mixture of Li2CO3 and Mn2O3 in air at 850° for 1 h. The use of λ-MnO2 in an organic-electrolyte Li

ES 1980-491736

US 1980-180616

19800522

19800825

battery was demonstrated.

ST manganese dioxide manuf battery; lithium battery

B4

A1

Α

manganese dioxide ΙT

JP 58034414

ES 491736

US 4312930

treatment

Batteries, primary

(lithium, organic-electrolyte)

TΤ Electrolytic depolarizers

(battery, manganese dioxide, manufacture of)

ΙT 1313-13-9P, preparation

RL: PREP (Preparation)

(manufacture of λ -, from lithium manganese oxide, by treatment with inorg. acid)

L97 ANSWER 7 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1967:121440 HCAPLUS

DN 66:121440

ED Entered STN: 12 May 1984

TΙ High energy electrochemical battery systems using organic electrolytes

ΑU Knapp, Howard R.

CS Army Electrons. Command, Fort Monmouth, NJ, USA

SO U. S. C. F. S. T. I., AD Rep. (1965), AD 627215, 55 pp. Avail.: CFSTI, \$3 СУ

From: U.S. Govt. Res. Develop. Rept. 1966, 41(5), 40

CODEN: XCCIAV

DT Report

LA English

CC 77 (Electrochemistry)

AB cf. CA 64, 3011d. Electrochemical studies on high energy anode and cathode materials coupled in organic electrolytes were undertaken to determine the feasibility of producing batteries superior to aqueous systems in

the areas of (1) higher energy ${\tt d}.$ per unit weight and volume, (2) wider temperature

range of operation than aqueous batteries, and (3) lower operating temperature

than

aqueous batteries. The report covers the selection of couples for study in organic electrolytes. Statistical evaluation of half-cell potential data at room temperature has disclosed inorg. acid anhydrides to be promising cathode materials in several organic electrolytes. Preliminary cell studies using Li anodes, dimethyl sulfoxide electrolytes, and acid anhydride cathode materials, such as Cro3, IZO5, molybdic oxide, and VZO5 were to characterize electrochem. performance of these materials.

ST ORG ELECTROLYTE BATTERIES; LITHIUM ANODES BATTERIES; BATTERIES ORG ELECTROLYTE

IT Organic compounds, uses and miscellaneous
RL: USES (Uses)

(electrolytes, in high-energy primary batteries)

IT Batteries, primary

(high-energy, with organic electrolytes)

=> => file wpix FILE 'WPIX' ENTERED AT 12:56:46 ON 03 JUN 2004 COPYRIGHT (C) 2004 THOMSON DERWENT

FILE LAST UPDATED: 3 JUN 2004 <20040603/UP>
MOST RECENT DERWENT UPDATE: 200435 <200435/DW>
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GUIDES, PLEASE VISIT:
http://thomsonderwent.com/support/userguides/

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>>> THE DISPLAY LAYOUT HAS BEEN CHANGED TO ACCOMODATE THE NEW FORMAT GERMAN PATENT APPLICATION AND PUBLICATION NUMBERS. SEE ALSO: http://www.stn-international.de/archive/stnews/news0104.pdf <<</p>

>>> SINCE THE FILE HAD NOT BEEN UPDATED BETWEEN APRIL 12-16 THERE WAS NO WEEKLY SDI RUN <<<

=> d que
L66 32608 SEA FILE=HCAPLUS ABB=ON (LI OR LITHIUM) (5A)BATTER?
L67 4 SEA FILE=HCAPLUS ABB=ON L66 AND ORG? AND INORG? (1W)ACID
L68 6 SEA FILE=HCAPLUS ABB=ON L66 AND ORG? (6A) INORG? (1W) ACID
L70 0 SEA FILE=HCAPLUS ABB=ON L66 AND ORG? (2A) BASE? AND INORG? (2A) AC
L71 1 SEA FILE=HCAPLUS ABB=ON L66 AND ORG? (2A) BASIC? AND INORG? (2A) A
CTD?
L98 11 SEA FILE=WPIX ABB=ON L67 OR L68 OR L70 OR L71

<<<

=> d 198 all 1-11

L98 ANSWER 1 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2004-106569 [11] WPIX

DNN N2004-084685 DNC C2004-043285

TI New carbon material carbonized/prepared from poly(phenylene ether) is used in adsorption applications, e.g. anti-pollution devices.

DC A18 A21 A25 A35 A85 D15 E36 J01 J06 L03 V01 X16

IN CABASSO, I; LI, S; LIU, H; YUAN, Y

PA (CABA-I) CABASSO I; (LISS-I) LI S; (LIUH-I) LIU H; (YUAN-I) YUAN Y

CYC 1

PI US 2003161781 A1 20030828 (200411)* 23 C01B031-02

ADT US 2003161781 A1 US 2001-968290 20011001

PRAI US 2001-968290 20011001 IC ICM C01B031-02

IC ICM C01B031-02 AB US2003161781 A UPAB: 20040213

NOVELTY - A carbon material carbonized/prepared from a poly(phenylene ether) and comprising 60-99.99 weight% carbon is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for:
(a) a composite comprising carbon and a modified poly(phenylene ether) (MPPE) carbon material, in which carbon material binds carbon particles and/or carbon fibers while maintaining the shape of the composite; and

(b) preparing an activated carbon from poly(phenylene ether) (PPE) precursor polymer of formula (I), by oxidizing the poly(phenylene ether) precursor, and carbonizing the modified poly(phenylene ether);

(c) preparing carbon/carbon composite from MPPE by forming a carbon fiber fabric with an organic fibrous binder consisting of cellulose, cellulose ethers or its derivatives, polyacrylonitrile, oxidized polyacrylonitrile, phenolic resins, polyvinyl acetate and/or epoxides; forming a solution of PPE or a slurry by mixing the PPE solution with other carbonous materials; applying the PPE solution to the carbon fiber fabric to form a composite; drying the composite; pressing the dried composite at 1-10000 psig; oxidizing the pressed composite in an oxygen containing atmosphere at 100-420 deg. C to crosslinked the PPE polymer to form MPPE; carbonizing the MPPE at 500-3000 deg. C in non-oxidizing atmosphere to form a carbon material; and activating the MPPE electrode with oxidizing gases, bases, or acids at elevated temperatures; and

```
(d) forming porous electrodes for double layer capacitors by forming
a slurry by mixing PPE solution with other carbonous materials;
transferring the slurry into a body having holes or dents on it; drying
the slurry to form a solid body; pressing the solid body at 10-5000 psig;
oxidizing the body in an oxygen containing atmosphere; carbonizing the
MPPE composite to form an MPPE electrode; activating the MPPE electrode;
and assembling a double layer capacitor with MPPE electrode by sandwiching
them with a separator in an electrolyte, and stacking with current
collectors.
```

R1, R2 = H, 1-6C aliphatic group, 6-24C aromatic group, aliphatic ether, aromatic ether, aliphatic or aromatic ester, carbonyl ester, carboxylic acid, ketone, lactone, or xanthone; n = 10-10000.

Intermolecular and intramolecular linkages exist between the R1 and R2 groups in the MPPE.

USE - Used in adsorption applications, e.g. anti-pollution devices, fuel gas storage, or electrochemical applications including electrodes or current collectors for double layer capacitors and lithium-ion batteries, or gas diffusion electrodes for fuel cells.

ADVANTAGE - The carbon material having high electric conductivity, high surface area controllable pore size distribution, long lifetime, low strength deterioration, and is easy to manufacture. It has a porosity of 10-90%, a maximum pore diameter of 0.00015-500 mu m, and BET surface are before activation of 500-700 m2/g.

DESCRIPTION OF DRAWING(S) - The figure is a labeled flow chart of the method for forming the carbon material/carbon composite. Dwg.1/21

FS CPI EPI

MC

FA AB; GI; DCN

CPI: A05-H06; A10-E05B; A12-E07B; D04-A01F; E31-N03; J01-D01; J01-E02B; J06-B06; L03-J

EPI: V01-B01A; V01-B01D; X16-E01C; X16-E01E; X16-E02; X16-E06A; X16-E08A; X16-L02

L98 ANSWER 2 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2003-681889 [65] WPIX

DNN N2003-544470 DNC C2003-186352

TI Porous hydrophilic membrane for fuel cell, comprises amorphous ionomer having acidic hydrophilic group deposited on porous inert support, and has ionic conductivity in electrochemical cell, and preset water permeability.

DC A14 A25 A85 L03 X16

IN ARCELLA, V; GHIELMI, A

PA (AUSY) AUSIMONT SPA

CYC 30 PT EP

EP 1238999 A1 20020911 (200365)* EN 14 C08J005-22

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

CA 2373396 A1 20020826 (200365) EN B01D071-36 CN 1371934 A 20021002 (200365) C08J005-18 JP 2002334707 A 20021122 (200365) 12 H01M008-02 US 2002144899 A1 20021010 (200365) C25B013-00

ADT EP 1238999 A1 EP 2002-3033 20020212; CA 2373396 A1 CA 2002-2373396 20020226; CN 1371934 A CN 2002-105353 20020226; JP 2002334707 A JP 2002-50044 20020226; US 2002144899 A1 US 2002-80555 20020225

PRAI IT 2001-MI383 20010226

IC ICM B01D071-36; C08J005-18; C08J005-22; C25B013-00; H01M008-02 ICS B01D053-22; B01D069-10; B01D071-32; B01D071-34; C08F214-18; C08J005-20; C08J009-42; C08L027-12; C25C007-04; H01B001-06; H01M008-10

ICI C08L027:18

AB EP 1238999 A UPAB: 20031009

NOVELTY - A porous hydrophilic membrane comprises an ionomer which is deposited on a porous inert support. The membrane has ionic conductivity in electrochemical cell and water permeability of more than 1 1/(h.m2.atm). The ionomer in amorphous form comprises acid form hydrophilic group.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for preparation of hydrophilic porous membrane which involves impregnating porous support in solution of ionomeric compound having fluorinated organic solvent at 15-40 deg. C to form membrane. The membrane is thermally treated at 120-160 deg. C to remove solvent, and to form transparent membrane. The membrane is treated with inorganic aqueous base alkali, and the functional groups are converted into hydrophilic groups. The obtained membrane is treated in aqueous inorganic strong acid to form (per)fluorinated ionomer in acid hydrophilic form. The obtained membrane is optionally treated with water at 50-100 deg. C to remove excess ionomer.

USE - Such as proton exchange membrane for electrochemical cells and fuel cells (claimed), and occluded membrane is used as semi-permeable membrane for reverse osmosis. The partially occluded porous membrane is used for forming electrodes and separators for lithium

ADVANTAGE - The hydrophilic membrane has high water permeability, high conductivity in electrolytic cells, and improved proton exchange capability. The membrane having ionomer of low equivalent weight, has superior application efficiency. The membrane when used in the catalytic reactions, shows high catalytic activity and when the membrane is used in the electrochemical applications, shows high ionic exchange capability. The membranes are used at room temperature, and high and low temperatures. Dwg.0/0

FS CPI EPI

FA AB

MC CPI: A04-E10; A10-E01; A11-B05; A12-E06B; L03-B01A; L03-E01B5; L03-E04 EPI: X16-B01F1; X16-C; X16-F02

L98 ANSWER 3 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2002-231328 [29] WPIX

DNN N2002-178031 DNC C2002-070452

TI Anode active material manufacture for non-aqueous electrolyte battery, involves immersing specific lithium complex into process liquid containing polar organic solvent and inorganic acid for specific time.

DC L03 X16

PA (TOYT) TOYOTA JIDOSHA KK

CYC 1

PI JP 2001332261 A 20011130 (200229)* 7 H01M004-58

ADT JP 2001332261 A JP 2000-150317 20000522

PRAI JP 2000-150317 20000522

IC ICM H01M004-58

ICS H01M010-40

AB JP2001332261 A UPAB: 20020508

NOVELTY - Lithium complex such as lithium nickelate of rock salt structure, is immersed into a process liquid containing a polar organic solvent and inorganic acid, for 12 hours or more.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for non-aqueous electrolyte battery.

USE - For non-aqueous electrolytic battery (claimed) such

as lithium-ion secondary battery.

ADVANTAGE - Non-aqueous electrolyte battery with excellent durability and with stabilized discharge characteristics, is obtained.

DESCRIPTION OF DRAWING(S) - The figure shows the explanatory and cross sectional views of change of crystal structure by immersion process. Dwg.2/6

FS CPT EPT

FΑ AB: GI

MC CPI: L03-E01C2

EPI: X16-B01F; X16-E01C; X16-J02; X16-J08

L98 ANSWER 4 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

2001-608965 [70] ΑN WPTX

DNN N2001-454759 DNC C2001-181244

Cation analysis involves introducing sample containing cation into cation exchange resin column, and separating and analyzing cation using aqueous solution containing inorganic acid and organic acid as mobile phase.

DC E37 J04 L03 S03

PA (SEIM-N) SEIMI CHEM KK

CYC 1

PI JP 2001174446 A 20010629 (200170) * 6 G01N030-88 ADT JP 2001174446 A JP 1999-360205 19991220

PRAI JP 1999-360205 19991220

IC ICM G01N030-88

AB JP2001174446 A UPAB: 20011129

> NOVELTY - The sample solution containing a cation is introduced into a cation exchange resin column of an ion chromatography apparatus. The cation in the sample is separated and analyzed using an aqueous solution which contains inorganic acid and an organic acid as a mobile phase.

USE - For analysis of cation in sample, for analysis of elementary composition ratio in positive electrode active material of lithium secondary battery.

ADVANTAGE - The quantity of alkali metal, alkaline earth metal and transition metal or measured simultaneously and accurately. The elementary composition ratio in positive electrode active material of lithium secondary battery is measured quickly and accurately. Analysis of semiconductor processing components, active materials and impurity element present in trace amount of fuel cell and nickel hydrogen battery, and lapping powder slurry for chemical mechanical polishing, are also enabled.

Dwg.0/1 FS CPI EPI

FA AB; DCN

MC CPI: E10-C02; E10-C04J1; E10-C04J2; E11-O03; E31-B03D; E31-F04; E31-H05; E33; E34; E35; J04-B01C; L03-E01B EPI: S03-E09C

L98 ANSWER 5 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2001-550252 [61] WPIX

CR 2000-349945 [30]; 2002-598993 [64]

DNC C2001-163866

TΙ Production of alkylene carbonate by contacting carbon dioxide, alkylene oxide, and catalyst to obtain crude effluent, subjecting effluent to multiple distillations, and further contacting purified alkylene carbonate with carbon.

DC E17

IN MACHAC, J R; MARQUIS, E T; SANDERSON, J R; SAVAGE, R L; WOODRUM, S A

(HUNT-N) HUNTSMAN PETROCHEMICAL CORP CYC 94 PΙ WO 2001066510 A2 20010913 (200161)* EN 26 C07C068-00 RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU ZA ZW AU 2001045460 A 20010917 (200204) C07C068-00 US 6384240 B1 20020507 (200235) C07D317-36 US 6458969 B1 20021001 (200268) C07D317-36 US 2002147349 A1 20021010 (200269) C07D317-08 WO 2001066510 A2 WO 2001-US7129 20010307; AU 2001045460 A AU 2001-45460 20010307; US 6384240 B1 CIP of US 1998-167361 19981007, US 2000-521502 20000309; US 6458969 B1 CIP of US 1998-167361 19981007, Div ex US 2000-521502 20000309, US 2002-71169 20020329; US 2002147349 A1 CIP of US 1998-167361 19981007, Div ex US 2000-521502 20000309, US 2002-71169 20020329 AU 2001045460 A Based on WO 2001066510; US 6458969 B1 CIP of US 6156160, Div ex US 6384240; US 2002147349 A1 CIP of US 6156160, Div ex US 6384240 PRAI US 2000-521502 20000309; US 1998-167361 19981007;

US 2002-71169 20020329 IC ICM C07C068-00; C07D317-08; C07D317-36

ICS B01D003-00; B01D003-34

AB WO 200166510 A UPAB: 20021026

NOVELTY - An alkylene carbonate is produced by contacting carbon dioxide, an alkylene oxide, and a carbonation catalyst in a reactor to produce a crude reactor effluent. The effluent is subjected to multiple distillations, where the unused fractions are recycled to the reactor. The purified alkylene carbonate is further contacted with carbon to reduce its ultraviolet absorbance.

DETAILED DESCRIPTION - Production of alkylene carbonate involves contacting carbon dioxide, alkylene oxide, and carbonation catalyst in a reactor to produce a crude reactor effluent. The effluent is subjected to low temperature evaporation to form an evaporator overhead containing alkylene carbonate and an evaporator bottom stream containing the catalyst. The evaporator bottom stream is recycled to the reactor, and the light components present in the evaporator overhead are removed to form a second evaporator overhead. The light components are recycled to the reactor. The second evaporator overhead is distilled to form a first distillation overhead stream and a first distillation bottom stream containing alkylene carbonate. The first distillation overhead stream is recycled to the reactor. The first distillation bottom stream is again distilled to form a second distillation overhead stream and a second distillation bottom stream, and the second distillation bottom stream is recycled. The second distillation overhead stream is further distilled to form a third distillation overhead stream and a third distillation bottom stream, and recycling the third distillation overhead stream. The third distillation bottom stream is further distilled to form a fourth distillation overhead stream containing purified alkylene carbonate and a fourth distillation bottom stream, and recycling the fourth distillation bottom stream. The purified alkylene carbonate is contacted with carbon to reduce its ultraviolet (UV) absorbance.

An INDEPENDENT CLAIM is also included for a process useful for reducing the UV absorbance of an alkylene carbonate.

USE - The method is for producing alkylene carbonate used as solvents for electrolyte salts in **lithium batteries**. The alkylene carbonate can also be used in electrochromic or photochromic

applications. ADVANTAGE - The inventive method generates less waste and is cost-effective. It produces alkylene carbonate of high purity (at least 99.99%) and very low ultraviolet absorbance (i.e., 0.350 at 220 nm). The alkylene carbonate has low water content (e.g., less than 20 ppm) and low glycol by-product (e.g., less than 20 ppm). Dwg.0/2 FS CPI FA AB; DCN MC CPI: E07-A04; N05-D L98 ANSWER 6 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN 2001-162029 [17] AN WPIX DNN N2001-118195 DNC C2001-048717 TI Recovery of cobalt from aqueous cobalt solution involves dissolving cobalt in organic acid, removing insoluble components and reducing cobalt by adding amphoteric metal. DC L03 M25 X16 PA (ASAK-N) ASAKA RIKEN KOGYO KK CYC 1 PΤ JP 2000328153 A 20001128 (200117)* 6 C22B023-00 ADT JP 2000328153 A JP 1999-133543 19990514 PRAI JP 1999-133543 19990514 IC ICM C22B023-00 ICS C22B003-04; C22B003-44; H01M010-54 ICA C22B007-00 AB JP2000328153 A UPAB: 20010328 NOVELTY - Cobalt is dissolved in an inorganic acid and insoluble components are removed from the solution by purification. To the reduced cobalt solution one or more amphoteric metal is added, cobalt is precipitated and recovered when the normal electrode potential of the reduction solution is in the range of -2.4 to -0.6 V. The amphoteric metal is then removed by alkali treatment. USE - For recovering cobalt from an aqueous cobalt solution obtained from lithium ion battery. ADVANTAGE - Cobalt is recovered efficiently. Dwg.1/2 FS CPI EPI FA AB; GI MC CPI: L03-E01; L03-E01C; L03-E03; M25-E01 EPI: X16-X L98 ANSWER 7 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN AN 2001-141944 [15] WPIX DNN N2001-103728 DNC C2001-042420 TТ Formation of solid polymer electrolyte for use in electrochemical devices involves reacting compound containing at least two sulfonic acid groups or its derivative with compound containing at least two amino groups. DC: A85 L03 X12 X16 PA (HITB) HITACHI CHEM CO LTD CYC 1 JP 2000331713 A 20001130 (200115)* PT 12 H01M010-40 ADT JP 2000331713 A JP 1999-141396 19990521 PRAI JP 1999-141396 19990521 IC ICM H01M010-40 ICS C08G075-30; C08K003-00; C08L081-10; H01B001-06; H01B013-00 AB JP2000331713 A UPAB: 20010317 NOVELTY - A compound containing at least two sulfonic acid, or its halide or other derivative is reacted with a compound containing at least two

amino groups. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following: (i) Solid polymer electrolyte; and (ii) electrochemical device using polymer electrolyte. USE - For formation of solid polymer electrolyte used for electrochemical device such as lithium secondary battery ADVANTAGE - The electrolyte has high ionic conductivity, less interfacial resistance, excellent toughness and mechanical properties. Electrochemical device with excellent thinning, laminate formation properties, easier packagability and reduced weight is obtained using the solid electrolyte. Dwg.0/0 FS CPI EPI FΑ AB MC CPI: A99-A; L03-E01C EPI: X12-D01C; X12-D07; X16-B01F L98 ANSWER 8 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN AN 1999-464225 [39] WPIX DNN N1999-347888 DNC C1999-136550 Electrolyte for lithium secondary battery - contains ΤI carbonate that acts as organic solvent and polymerisation inhibitor. DC. L03 X16 PA (FUIT) FUJITSU LTD CYC 1 PT JP 11195427 A 19990721 (199939)* 6 H01M010-40 ADT JP 11195427 A JP 1998-1070 19980106 PRAI JP 1998-1070 19980106 TC ICM H01M010-40 AB JP 11195427 A UPAB: 19991026 NOVELTY - The electrolyte (5) contains lithium salt with inorganic or organic acid residues excluding As. Carbonate as an organic solvent and polymerisation inhibitor is also included. USE - For lithium secondary battery. ADVANTAGE - Raise of temperature is suppressed and charging/discharging characteristics are improved. The flow of electricity is reduced as internal resistance is increased. Dwg.0/2 FS CPI EPI FA AB CPI: L03-E01C EPI: X16-B01F1; X16-J L98 ANSWER 9 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN 1998-600320 [51] AN WPIX DNN N1998-467710 DNC C1998-179899 Electrolyte for lithium sec. battery - uses a combination of an organic acid lithium salt, and at least one inorganic acid lithium salt. E12 L03 X16 (MITU) MITSUBISHI CHEM CORP CYC 1 JP 10270078 A 19981009 (199851)* 7 'H01M010-40 ADT JP 10270078 A JP 1997-75534 19970327 PRAI JP 1997-75534 19970327 ICM H01M010-40 JP 10270078 A UPAB: 19981223 An electrolyte uses a combination of an organic acid lithium

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salt of formula LiN(SO2CnF2n+1) (SO2CmF2m+1) (1), and at least one inorganic acid lithium salt selected from the group consisting of LiX (X = PF6, AsF6, SbF6, or BF4) with mol ratios ranging from 95:5 to 60:40 as a solute, and contains an ester carbonate cpd. as an organic solvent. n and m = 1-4.

ADVANTAGE - The solute has an oxidation electric potential for

ADVANTAGE - The solute has an oxidation electric potential for aluminium higher than an electric potential for a positive electrode in normal charge. The result exerts good charge with evolving less dissolution of the aluminium current collector of the positive electrode to the electrolyte. The electrolyte has superior conductivity, provides the aluminium of the current collector of the positive electrode in charge with no corrosion (no dissolution) to yield high lithium charge-discharge efficiency, resulting in good charge-discharge characteristics. High Dwg.0/1

FS CPI EPI

FA AB; DCN

MC CPI: E05-A; E11-N; L03-E01C

EPI: X16-B01F1; X16-J

L98 ANSWER 10 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN

AN 1998-527099 [45] WPIX

DNN N1998-412132 DNC C1998-158365

TI Producing anode active material for secondary battery - by adding organic amine to solution of lithium and other metal element, adding hydrogen peroxide to form hydroxide, mixing obtained powder with solution of composite fatty acid metal salt, etc.

DC E19 L03 X16

PA (MURA) MURATA MFG CO LTD

CYC 1

PI JP 10233213 A 19980902 (199845)* 5 H01M004-58 JP 3301335 B2 20020715 (200253) 5 H01M004-58

ADT JP 10233213 A JP 1997-32028 19970217; JP 3301335 B2 JP 1997-32028 19970217

FDT JP 3301335 B2 Previous Publ. JP 10233213 PRAI JP 1997-32028 19970217

IC ICM H01M004-58

ICS C04B035-495; H01M004-02; H01M004-04; H01M010-40

AB JP 10233213 A UPAB: 19981111

An anode active material consists of LiMxO2x. M = at least element selected from Ni, Mn, Co, Cr, Fe, and V; x = 1 or 2. The production comprises:

(a) adding an organic amine to a mixed solution containing Li+ and M (n = number of ion valencies of M element) to precipitate M(OH)n (n = number of ion valencies of M element); (b) adding hydrogen peroxide to the developed M(OH)n solution to form M(OH)n to MOOH and to deposit Li2O2 on the surfaces of the MOOH particles; drying the Li2O2-deposited MOOH particles; (c) mixing the MOOH powder with the solution of a composite fatty acid metal salt cpd. of Li and M; (d) drying the resulting mixture to cover the surface of the powder by the composite fatty acid metal salt cpd.; (e) heat treating the covered powder.

ADVANTAGE - The method synthesizes the homogeneous active material at low temps. The particle dia. is fine submicron. The battery has enhanced filling density of the active material to increase electrical capacity. An inexpensive inorganic acid salt is used as a starting material. No crushing process is needed, producing the active material at lower cost

lower cost. Dwg.0/0

FS CPI EPI FA AB; DCN

MC CPI: E10-B04; E10-C04L; L03-E01B8

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EPI: X16-B01F; X16-E01; X16-E01C; X16-E01G
L98 ANSWER 11 OF 11 WPIX COPYRIGHT 2004 THOMSON DERWENT on STN
      1993-296675 [38]
                        WPIX
DNN N1993-228676
                        DNC C1993-131456
тT
     Lithium-intercalating carbon electrode for storage
     battery - obtd. by heating organic or inorganic
     acid with carbon precursor to gives large capacity and freedom
      from dendrites.
DC
     A85 E17 E36 L03 X16
IN
     BITO, Y; HASEGAWA, M; ITO, S; MURAI, H; TOYOGUCHI, Y
PA
      (MATU) MATSUSHITA ELEC IND CO LTD; (MATU) MATSUSHITA ELECTRIC IND CO LTD
CYC 5
PΙ
     EP 561372
                     A1 19930922 (199338)* EN
                                                15
                                                      H01M004-58
         R: DE FR GB
     JP 06044959 A 19940218 (199412)
                                                11
                                                      H01M004-02
                    A 19961203 (199703)
     US 5580538
                                                9
                                                      D01F009-12
     EP 561372
                    B1 19970604 (199727) EN
                                               15
                                                      H01M004-58
         R: DE FR GB
     DE 69311170
                  E 19970710 (199733)
                                                      H01M004-58
     JP 3401646
                    B2 20030428 (200330)
                                                11
                                                      H01M004-58
ADT EP 561372 A1 EP 1993-104339 19930317; JP 06044959 A JP 1993-46528
     19930308; US 5580538 A Cont of US 1993-31627 19930315, US 1994-306248
     19940914; EP 561372 B1 EP 1993-104339 19930317; DE 69311170 E DE
     1993-611170 19930317, EP 1993-104339 19930317; JP 3401646 B2 JP 1993-46528
     19930308
FDT DE 69311170 E Based on EP 561372; JP 3401646 B2 Previous Publ. JP 06044959
PRAI JP 1992-61661
                         19920318; JP 1992-61665
                                                         19920318
REP 1.Jnl.Ref; EP 418514; EP 529095; JP 03245458; JP 3245458
IC
     ICM D01F009-12; H01M004-02; H01M004-58
     ICS C01B031-02; H01M004-04; H01M010-40
AB
     EP
           561372 A UPAB: 19931123
     A Li-intercalating negative electrode (1) for a storage battery including
     a reversible anode (4), a non-aqueous Li salt-containing electrolyte, and the
     cathode which is made of a carbon material having an improved surface area
     large enough to store Li ions to provide the largest possible charge and
     discharge capacitances.
          Also claimed is production of the cathode by mixing an acid with the
     C-containing material, heating to form the C material, and forming the
     electrode.
          Pref. the carbon material can form an intercalation cpd. with Li and
     is graphite. The acid is inorganic, sulphuric, nitric, or hydrochloric, or
     boric, or carboxylic such as acetic or formic. The C-containing material is
     organic- pitch, coal tar, coke, resin, cellulose, PAn, or rayon,
     or inorganic- graphite, carbon fibre, or graphite whisker. The
     organic material is heated at 400-3000 deg.C or where it is
     converted to inorganic C, and the inorganic material is heated at 100-1500
     deg.C.
          USE/ADVANTAGE - The electrode is useful for high energy density
     batteries, is reliable, increases charge-discharge capacity, and avoids
     shorts caused by dendrite formation.
     10
     Dwg.1/4
     CPI EPI
FS
FA
     AB; GI; DCN
MC
     CPI: A12-E06A; E31-N02; E31-N03; L03-E01B5
     EPI: X16-E01A; X16-E08A
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